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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
09/544,108	04/06/2000	Kenneth Eliot Sherman		7634
	7590 04/15/200 taff Judge Advocate	EXAMINER		
U.S. Army Medical Research and Material Command ATTN: MCMR-ZA-J (Ms. Elizabeth Arwine) 504 Scott Street Fort Detrick, MD 21702-5012			BOESEN, AGNIESZKA	
			ART UNIT	PAPER NUMBER
			1648	
			MAIL DATE	DELIVERY MODE
			04/15/2008	PAPER

## Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

## Advisory Action Before the Filing of an Appeal Brief

Application No.	Applicant(s)		
09/544,108	SHERMAN, KENNETH ELIOT		
Examiner	Art Unit		
Agnieszka Boesen	1648		

	Agnieszka Boesen	1648	
The MAILING DATE of this communication appe	ars on the cover sheet with the c	correspondence add	ress
THE REPLY FILED 11 February 2008 FAILS TO PLACE THIS	APPLICATION IN CONDITION FO	R ALLOWANCE.	
1. The reply was filed after a final rejection, but prior to or on application, applicant must timely file one of the following application in condition for allowance; (2) a Notice of Apperfor Continued Examination (RCE) in compliance with 37 C periods:	replies: (1) an amendment, affidavit eal (with appeal fee) in compliance	t, or other evidence, w with 37 CFR 41.31; or	hich places the (3) a Request
a) The period for reply expires 3 months from the mailing date b) The period for reply expires on: (1) the mailing date of this A no event, however, will the statutory period for reply expire to Examiner Note: If box 1 is checked, check either box (a) or ( MONTHS OF THE FINAL REJECTION. See MPEP 706.07(f)	dvisory Action, or (2) the date set forth in ater than SIX MONTHS from the mailing b). ONLY CHECK BOX (b) WHEN THE	g date of the final rejection	n.
Extensions of time may be obtained under 37 CFR 1.136(a). The date of have been filed is the date for purposes of determining the period of extunder 37 CFR 1.17(a) is calculated from: (1) the expiration date of the set forth in (b) above, if checked. Any reply received by the Office later may reduce any earned patent term adjustment. See 37 CFR 1.704(b). NOTICE OF APPEAL	ension and the corresponding amount of hortened statutory period for reply original controls.	of the fee. The appropria nally set in the final Offic	ate extension fee e action; or (2) as
<ol> <li>The Notice of Appeal was filed on A brief in comp filing the Notice of Appeal (37 CFR 41.37(a)), or any exter Notice of Appeal has been filed, any reply must be filed wi AMENDMENTS</li> </ol>	nsion thereof (37 CFR 41.37(e)), to	avoid dismissal of the	
<del></del>			
<ol> <li>The proposed amendment(s) filed after a final rejection, k</li> <li>(a) They raise new issues that would require further cor</li> <li>(b) They raise the issue of new matter (see NOTE below</li> </ol>	nsideration and/or search (see NOT		cause
(c) They are not deemed to place the application in bett appeal; and/or	ter form for appeal by materially rec	ducing or simplifying t	ne issues for
(d) They present additional claims without canceling a converse NOTE: (See 37 CFR 1.116 and 41.33(a)).	corresponding number of finally reje	ected claims.	
4. The amendments are not in compliance with 37 CFR 1.12	21 See attached Notice of Non Co.	mpliant Amandment (	DTOL 224\
5. Applicant's reply has overcome the following rejection(s):		mpliant Amendment (i	-10L-324).
<ol> <li>Newly proposed or amended claim(s) would be all non-allowable claim(s).</li> </ol>	owable if submitted in a separate, t	imely filed amendmer	nt canceling the
7. For purposes of appeal, the proposed amendment(s): a) how the new or amended claims would be rejected is prove The status of the claim(s) is (or will be) as follows:		l be entered and an e	xplanation of
Claim(s) allowed: Claim(s) objected to: Claim(s) rejected: <u>1,3,4,6 and 25</u> . Claim(s) withdrawn from consideration:			
AFFIDAVIT OR OTHER EVIDENCE			
<ol> <li>The affidavit or other evidence filed after a final action, but because applicant failed to provide a showing of good and was not earlier presented. See 37 CFR 1.116(e).</li> </ol>			
<ol> <li>The affidavit or other evidence filed after the date of filing entered because the affidavit or other evidence failed to o showing a good and sufficient reasons why it is necessary</li> <li>The affidavit or other evidence is entered. An explanation</li> </ol>	vercome <u>all</u> rejections under appea and was not earlier presented. Se	al and/or appellant fail ee 37 CFR 41.33(d)(1	s to provide a ).
REQUEST FOR RECONSIDERATION/OTHER	TOT THE Status OF THE Claims after er	illy is below of allacin	cu.
11. The request for reconsideration has been considered but	does NOT place the application in	condition for allowan	ce because:
12. ☐ Note the attached Information <i>Disclosure Statement</i> (s). ( 13. ☐ Other:	PTO/SB/08) Paper No(s)		
	/Agnieszka Boesen, Ph Examiner, Art Unit 1648	n.D./	
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## **Continuation Sheet (PTO-303)**

Application No.

Applicant's arguments have been fully considered but fail to persuade. Applicant argues that in the Declaration by Dr. Kenneth Sherman submitted on January 2, 2006, Applicant's have shown improved results achieved with the interferon alpha and thymosin alpha combination therapy over using interferon alpha or thymosin alpha alone. The Declaration by Dr. Sharman has been considered. It is noted that Figure 2 shows the results of treatment with the combination therapy and with interferon alpha alone. It is acknowledged that the combination therapy resulted in higher histologic response in the HCV patients as compared to the treatment with interferon alpha alone. However this result with combination therapy would have been expected at the time of the present invention, because the skilled artisan had known that thymosin alpha acts on interferon alpha to increase the effects of interferon alpha regardless the type of virus that infects the host. Thus it has been known that thymosin alpha acts to increase the effects of interferon alpha, and therefore one would have expected that combination therapy would have given better results over interferon alpha alone in treatment of viral liver infection. The fact that interferon alpha was beneficial in treatment of both HBV and HCV infection has been known at the time of the present invention and as discussed on the record in the Office Action of 5/31/2007. Thus interferon alpha acts regardless the type of viral infection. Thymosin alpha acts on interferon alpha acts both on HBV and HCV. Thus because it has been known that interferon alpha acts both on HBV and HCV the present invention would have been obvious to the skilled artisan at the time when the invention was made. Therefore the rejection is maintained.

/Stacy B. Chen/ 4-10-2008 Primary Examiner, TC1600